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(54) Title: ACRYLIC BASED POWDER PAINT

(57) Abstract

The invention relates to a powder paint based on acrylics. The thermosetting powder paint comprises a hydroxyl functional acrylic resin and an isocyanate crosslinking agent blocked with a blocking agent wherein the isocyanate crosslinking agent is an adduct of an isocyanate and a mixture of polyetramethylene ether glycol and at least one additional, but other glycol, selected from the group consisting of glycols and polyglycols.

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ACRYLIC BASED POWDER PAINT

5 The invention relates to a powder paint based on acrylics.

The mechanical properties of acrylic based powder coatings are worse (pages 162-167 of "Powder Coatings, Chemistry and Technology" by Tosko Misev; John Wiley and Sons, 1991). Acrylic based powder coatings, 10 which are used in practice, are almost exclusively based on epoxy functional acrylic resins cured with long-chain dicarboxylic acids as hardeners. The long aliphatic chain of the crosslinker provides flexibility and impact 15 resistance to the cured film, but still far below the values which are usually obtainable with the other powder coating systems. In most cases the impact resistance of the acrylic powder coatings does not exceed 30 inchpounds. This is still several times lower compared to the 20 values of polyester and polyurethane based powder coatings.

It is the object of the invention to provide an acrylic powder coating system which shows excellent hardness, gloss, impact resistance, durability, clearness, chemical resistance and good flexibility.

The invention is characterized in that the thermosetting powder paint comprises

- a hydroxyl functional acrylic resin and

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- an isocyanate crosslinking agent blocked with a blocking 30 agent, wherein the isocyanate crosslinking agent is an adduct of an isocyanate and a mixture of polytetramethylene ether glycol and at least one additional, but other glycol, selected from the group consisting of glycols and polyglycols.
- 35 Preferably the crosslinker has a melting point high r than 45°C and a molecular weight M, between 100 and 3000, more preferably between 300 and 1500.

The mixture of glycols and/or polyglycols

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contains pr ferably more than 50% by w ight of polytetramethylene ether glycol as one polyol. More preferably this amount is between 60 and 75% by weight.

Suitable other glycols or polyglycols include a great number of diols, triols and polyols, such as for example butanediol, ethanediol, neopentylglycol, 2,2-bis-(1-hydroxy-2-oxyethylphenyl)propane, 1,1-isopropylidene-bis(phenylene-oxy)di-2-propanol-2, cyclohexyldimethylol, trimethylolpropane, trimethylolethane, adipate polyols, polycaprolactone glycols and polycarbonate glycols.

According to a preferred embodiment of the invention the second additional polyglycol is a polyether polyol. A suitable polyether polyol is Voranol 220-530TM (of Dow).

A suitable polytetramethylene ether glycol is Terathane 1000TM (of Du Pont).

Suitable isocyanates include aliphatic (poly)isocyanates such as for example hydrogenated methylene diphenyldiisocyanate (HMDI), isophorone

20 diisocyanate (IPDI), trimer(isocyanate) of isophorone dissocyanate (T1890TM, Hüls), 1,6-hexamethylene diisocyanate (HDI) and the trimer of 1,6-hexanediisocyanate (Tolonate HDTTM, Rhone Poulenc), 1,3-bis-(1-isocyanato-1-methylethyl)-benzene (TMXDI; American Cyanamid) or aromatic polyisocyanates, such as for example 2,4 or 2,6-diisocyanatetolueen (TDI) and 4,4'-di-isocyanatediphenylmethaan.

Preferably HMDI is used.

The flexible crosslinker can be prepared by

30 first reacting half of the molar equivalents of an
isocyanate with the polyglycol mixture and next reacting
the remaining isocyanate equivalent with a blocking agent.

A blocked isocyanate is an isocyanate which has been reacted with a mat rial which will prevent its reaction at roomtemp ratur with compounds that conventionally react with isocyanates but will p rmit that r action to occur at high r temp ratur.

Blocked isocyanates are described by Wicks in

Progress in Organic Coatings (3, 1975, 73-99).

According to a pref rr d embodim nt of th invention the blocking agent is a non-volatile agent having a polymerisable double bond and an oxime function.

These agents have the general formula (I):

$$R^{1}$$
 R^{3}
 $C=C-R^{4}-C-R^{5}$
 N
OH

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where

 R^1 , R^2 , R^3 = H or (C_1-C_5) alkyl

 R^4 = aliphatic, amidegroup or organic estergroup and 15 R^5 = (C_1-C_5) alkyl.

Preferably the amidegroup is steric hindered.

The aliphatic group, preferably, contains 1-10 carbon atoms.

The organic estergroup can be characterized by

20 0 $-C-O-R^6$ in which $R^6=(C_1-C_{10})$ alkyl.

The agent having a polymerizable double bond and an oxime function is preferably an acrylamide derivative having the formula (II):

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$$CH_{2} = C - C - N - C - CH_{2} - C - CH_{3}$$

$$O H CH_{3}$$

$$OH$$
(II)

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This suitable blocking agent according to the invention is diacetoneacrylamide oxime (DAAOX). The preparation of DAAOX is disclosed in Macromolecules, Vol. 16, 10, 1983, pag s 1561-1563.

The use of DAAOX and other blocking agents having a polymerisable double bond and an oxime function is very advantageous, because these blocking agents do not evolve during the cure of a hydroxy-isocyanate powder

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paint system. The blocking agent can be polymerised during cure of the powder paint, because they consist of a blocking side and a polymerizable double bond. The polymerizable double bond can be polymerized during the cure of the coating by adding a suitable peroxide. Any peroxide having a suitable halflife-time at the curing temperature of the coatings can be used. Preferably the peroxide has a halflife-time of less then 10 minutes at the temperature at which the coating is cured. The amount of peroxide can vary between 0.5 and 4 wt.%, preferably between 1 and 2%, based on the weight of the added blocked isocyanate crosslinker. Suitable peroxides include tert.—amyl-peroxybenzoate, tert.—butylperoxybenzoate and tert.—butylperoxy-2-ethylhexyl carbonate.

obtained from compounds having a ketone or aldehyde group which can be converted into an oxime and having an unsaturated double bond as well. Suitable examples of unsaturated ketones or aldehydegroups include

methyl-vinyl-ketone, ethyl-vinyl-ketone, mesityl-oxide, allylacetone, crotonaldehyde, 2-hexanal and citronellal. The blocking agent can be reacted with a (poly)isocyanate in a conventional way. Preferably the molar equivalent ratio (poly)isocyanate: blocking agent is substantially 1:1.

The main features of the isocyanate-hydroxyl curing reaction are described in the foregoing cited Powder Coatings, Chemistry and Technology at pages 56-58.

The use of a non-volatile agent having a polymerizable double bond and an oxime function is not limited to the reaction between a blocked isocyanate and a hydroxyl functional acrylic resin. The hydroxyl functional resin can also be, for example, a polyester or a polyurethane.

Suitable volatile blocking agents includ for example phenol, cresol, long-aliphatic-chain substitut d phenols such as isononylphenol, amid s such as \(\epsilon\)-caprolactam, activ methylen group containing compounds

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like malonates such as isopropyliden malonate and acetoacetic sters, sodium bisulfite and oximes such as for exampl methylethylketone oxime and butanone oxime.

Suitable hydroxyl-functional acrylate resin include for example resins based on hydroxyethyl (meth)acrylate, hydroxypropyl (meth)acrylate and methyl (meth)acrylate. The resin may also be based on methacrylic acid and alkyl esters of (meth)acrylic acid such as methyl acrylate, ethyl (meth)acrylate, isopropyl (meth)acrylate, n-butyl (meth)acrylate, n-propyl (meth)acrylate, isobutyl (meth)acrylate, ethylhexyl acrylate and/or cyclohexyl (meth)acrylate and vinyl compounds such as styrene.

Preferably, the hydroxylacrylate resins have a hydroxyl number between 40 and 150 mg KOH/g resin and an acid number lower than 20 mg KOH/g resin.

The acrylate resins can be prepared by a (co)polymerization, where solvent is fed to a reactor and then heated until the solvent boils. Monomers, and subsequently monomers, initiator and, optionally, mercaptan are added during a period of, for instance,

- mercaptan are added during a period of, for instance, between 2 and 4 hours, after which the temperature is kept at the reflux temperature for, for instance, two hours. The solvent is distilled off by increasing the temperature followed by a vacuum distillation lasting, for instance,
- one to two hours. Then the product is drained and cooled. Suitable solvents include for example toluene, xylene and butylacetate. Suitable initiators include azo-bis-isobutyronitrile, dibenzoylperoxide and tert.-amyl-peroxy-2-ethyl-hexanoate.
- The weight ratio polymer:crosslinker is generally between 90:10 and 50:50.

Preferably the molar equivalent ratio of polymer:crosslinker is between 1:0,8 and 1:1,5. Very suitable ratios are between 1:1 and 1:1,2.

Obviously, all customary additives may be included in the composition. Examples of customary additives include pigments, fillers, flow aids, stabilizers and catalysts. Suitable pigments include

inorganic pigments, for instanc titanium oxide, zinc sulphide, iron oxide and chromium oxide, and organic pigments, for instance azo compounds. Suitabl fill rs include metal oxides, silicates, carbonates and sulphates.

The technology and production of powder coatings is described at pages 224-226 of Powder Coatings, Chemistry and Technology, by Tosko Misev (1991; John Wiley and Sons).

Powder coatings according to the present

invention can be applied in general industrial and
domestic appliances, metal furnitures, architectural
applications, automotive topfinishes, corrosion protective
coatings and finishes for wood, plastics and paper.

US-A-5097010 relates to the preparation of 15 thermally-reversible isocyanate polymer by reacting a labile hydrogen segment with an isocyanate segment. The obtained compositions are useful as hot-melt adhesives, coatings and mouldings and furthermore in injection reaction moulding applications and composite and laminate 20 manufacturing followed by thermal forming and pulltrusion. In contrast to the present invention, it is the object of US-A-5097010 to provide a thermally-reversible system. Futhermore in contrast to the present invention, which relates to thermosetting coatings, said US-patent 25 discloses a thermoplastic system with polymer networks which are insoluble strong solids at room temperature but become soluble free flowing melts at high temperature. Of course the polyurethanes are based on disocyanates and polyols and a man skilled in the art also knows that long flexible polyols result in flexible polymers. The present 30 invention, however, relates to a flexible network and does not relate only to a flexible resin. It is for a man skilled in the art not obvious that a flexible part in a r sin will also r sult in a flexible network. It is th essential f ature of th pr sent inv ntion that the poly-35 tetramethyl ne ether glycol incorp rat d in a polyurethane resin (tog th r with oth r diols and diisocyanates) in combination with an acrylate r sin results in

a flexibl network. Applying polytetramethylen th r glycol as the glycol alone will not result in a superior powder coating resin. The mixture of polytetramethyl ne ether glycol with at least one other glycol or polyglycol produces the desired properties. One glycol is responsible for the flexibility, the other glycol is responsible for the level of the glass transition temperature.

The coatings of the present invention are further illustrated by the following experiments and examples. The examples are included for illustrative purposes and should not be considered to limit the present invention.

Experiment I

15 Synthesis of an hydroxyl functional acrylic resin

A 2 L reactor vessel, equipped with a thermometer, a stirrer and a reflux cooler, was charged with 500 g toluene. The reactor was stirred, a nitrogen flow passed through the reactor and the temperature was increased to reflux temperature. A monomer mixture consisting of 417 g methyl methacrylate (MMA), 175 g hydroxyethyl methacrylate (HEMA), 200 g cyclohexylmethacrylate (CHMA), 200 g n-butylacrylate (BA) and 8 g methacrylic acid (MA) was added. In this monomer mixture 29,4 g 2,2-azo-bis-isobutyronitrile (AIBN) was dissolved.

The monomer mixture was fed to the reactor in 2.5 hours. The reflux temperature was maintained in the reactor for another 2.5 hours. Then a separator vessel was included in the setup, and the solvent was removed by a gradual increase of the temperature and application of a vacuum. A clear product was obtained with a glass transition temperature (Tg, Mettler TA-3000, system 5°C/min) of 46°C and a viscosity (h) (measured with Emila rheometer, 165°C) of 350 dPa.s.

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Experiment II

Synthesis of diaceton acrylamid oxime (DAAOX)

100 g diac ton acrylamide was dissolved in 375

ml of distilled wat r and 45.75 g hydroxylamine hydrochlorid was added with stirring. A solution of 43.75 g potassium carbonat in 62.5 ml of distilled water was added at room temperature over a period of 20 min. After 5 the addition was completed, most of the oxime product precipitated. The reaction was continued for an additional hour and the precipitate was filtered and washed with 50 ml of ice water twice. The wet crude product was dissolved in ethylacetate (500 ml), the organic layer was separated and crystallized in the refrigerator. The crystallized product was filtered and washed with 100 ml hexane twice. The final product was dried at 40°C in a vacuum oven to yield 77 g of the DAAOX product. The melting point (112°C) of DAAOX was measured by using a DSC.

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Experiment III

The blocking of IPDI with DAAOX (BIPDI)

40.48 g (10% excess) DAAOX obtained according to Experiment I was dissolved in 50 g methylethyl ketone and 20 22.2 g IPDI was added. The reaction was carried out at room temperature under dry nitrogen. The NCO content was less then 0.1% (by titration) after one hour reaction. The solvent was removed by destillation at 40°C in a vacuum oven overnight. The final blocked IPDI (BIPDI) can be pulverized into fine powder with a melting point of 55-60°C. The blocked NCO content of BIPDI is 13.5% by calculation.

Experiment IV

30 The blocking of H12MDI with DAAOX (BHMDI)

147.5 g DAAOX (5% excess) obtained according to Experiment I was dissolved in methylethyl ketone (330 g) and 100 g H₁₂MDI was then added. The reaction was carried out at 50°C under dry nitrogen. Th NCO content was less then 0.1% (by titration) aft r one hour of reaction. The solv nt was r moved at 40 C in a vacuum ov n. The final block d product BHMDI was pulverized to a fine powder and had a melting point (by DSC) of appr. 54°C.

In order to xamine th polym rization of BHMDI, the intiator (tert.-amylperb nzoate) was add d into BHMDI to initiate polym rization. An exothermic peak was found in the analysis of DSC.

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Examples I-VII

Preparation of modified crosslinkers

The glycol blend was made by mixing two glycols in a ratio as mentioned in Table I. In a reactor, equipped with a stirrer and under a stream of dry nitrogen the 10 respective glycol blend and the appropriate amount of HMDI (see Table I) were charged. Methylethylketone (MEK) was added to reduce the viscosity. A small amount of dibutyltindilaurate was added (see Table I) as a catalyst, the reaction mixture was heated to 70°C. It was kept at that 15 temperature until the NCO percentage (as determined by titration) was reduced to half of the initial value. DAAOX was dissolved in MEK and then added into the reaction mixture. The reaction temperature was kept at 70°C until the NCO peak (2270 cm⁻¹) disappeared by checking the 20 infrared spectrum of the product.

After the products in MEK were further diluted with acetone to 25-30% solids content. The dilute solutions were then added dropwise into cold water under vigorous agitation (> 1500 RPM). The precipitates in the powdered form were then filtered and dried. Glass transition temperature and equivalent weights of NCO are given in Table I.

30 Examples VIII-XIV

Preparation of a powder coating

The powder coating ingredients were weighted (in grammes) in the formulations as showed in Table II. The formulations were dissolved is ac ton to prepar a solution of 25-30% solids cont nt. The solutions w r added into cold water dropwise und r vigorous agitation. The precipitat s in powd r form were then filt r d and dried. The dried powders were ground into a v ry fin

particle siz . They w re sprayed el ctrostatically on plates and cured as showed in Tabl II.

The present tests are described at pages 296-303 of Powder Coatings by Misev.

Comparative Example A

0.942 g of BHMDI according to Experiment IV was mixed with 2.058 g acrylic resin according to Example I, 10 and the powder mixture was dissolved in 7.5 g of cellosolve acetate. 0.0015 g dibutyltin diacetate was added as a deblocking catalyst. 0.0188 t-amyl-perbenzoate was added as initiator for the polymerization of DAAOX after deblocking reaction. 0.0585 g of E-25TH was added as flow control agent. The solution of this mixture was 15 coated on stell panels, and the panels were baked at 160°C for 30 min. The coatings were well cured, which was proved by the pencil hardness of 2H and more then 200 aceton rubs. However, the impact resistance (ASTM-D-2794) was 20 less than 30 inchpound.

This shows that the coating made with a non-modified crosslinker has poor properties.

The examples show that the isocyanate crosslinker according to the invention results in very flexible acrylic powder coatings with good properties. The claimed invention results in a remarkable increase in impact resistance.

TABLE I (grammes)

ABLE I

ស		·	VIII	IX	×	XI	XII	XIII	XIV
	R sin Exp. I		100	100	100	100	100	100	100
	Crosslinker I		105,3		ı	1) 	, , ,)) !
•	_		:	105,8	. 1	ı	ı		ı
	_		ı	. 1	96,85	. 1	ı	ŧ	
10	Crosslinker IV		ı	ı		111,0	1	ı	ı
	Crosslinker V		1	1		!	113,1	:	
		<i>:</i>	ı		ı	1		80,61	i .
	Crosslinker VII		f	ı	1	1	1		82,82
(TIT)	,	•	•	96′	, 11	2,131	1,806	82
15	•		1,053	1,058	0,9685	1,110	1,131	908'0	
	E(2 3)		•	•	, 55	,74	2,770	2,348	, 37
	Cur conditions		150.0	150°C	150.0	150°C	150°C	150°C	150.0
			30,	30,	30,	30,	30,		
20	Hardness ⁴⁾		H ^	H <	2H	H ^	H ^	H ^	H ^
	Aceton res. 5)		> 200	> 200	> 200	> 200		> 200	
	Impact res: 6)		•						
	- direct		120	110	110	-	16	> 160	140
7	- reverse		100	09	09 >	> 160	> 160	> 160	
0 0	TI ¹⁾ TAP ²⁾ E ^{25 3)} Hardness ⁴⁾ Ac ton res. ⁵⁾ Impact res. ⁶⁾	= Dibutyl = Tert. a = Flow co = Pencil = Acetone = Impact	ltin amyl ontro haro e res	eta enz ent (A	M) D-3363 Ivent M-D-27) cure test 94)	method)		

CLAIMS

- 1. A powder paint bas d on acrylics, charact riz d in that, the thermosetting powder paint comprises

 a hydroxyl functional acrylic resin and
 an isocyanate crosslinking agent blocked with a blocking agent, wherein the isocyanate crosslinking agent is an adduct of an isocyanate and a mixture of polytetramethylene ether glycol and at least one additional, but other glycol, selected from the group consisting of glycols and polyglycols.
 - 2. Powder paint according to claim 1, characterized in that the additional polyglycol is a polyether polyol.
- 3. Powder paint according to any one of claims 1-2,
 characterized in that the blocking agent is an agent
 having a polymerisable double bond and an oximefunction.
 - 4. Powder paint according to claim 3, characterized in that the agent is diacetone acrylamide oxime.
- 20 5. Use of an adduct of an isocyanate and a mixture of polytetramethylene ether glycol and at least one additional but other glycol selected from the group consisting of glycols and polyglycols as crosslinking agent in the preparation of powder coatings.
- 25 6. A method for preparing a powder paint composition comprising a hydroxyl functional acrylic resin and an isocyanate crosslinking agent which is blocked, characterized in that the isocyanate crosslinking agent comprises an adduct of an isocyanate and a mixture of polytetramethylene ether glycol and at least one additional, but other glycol selected from the group consisting of glycols and polyglycols and the weight ratio of polymer:crosslinking agent is between 90:10 and 50:50.
- 35 7. Method according to claim 6, charact rized in that the blocking ag nt is an agent having a polymerisable double bond and an oxim -function.

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- 8. Method according to claim 7, characterized in that th agent is diac ton acrylamide oxime.
- 9. Binder composition comprising a resin and a crosslinking agent characterized in that the resin is a hydroxyl functional acrylic resin and the crosslinking agent an isocyanate crosslinking agent blocked with a blocking agent, wherein the isocyanate crosslinking agent is an adduct of an isocyanate and a mixture of polytetramethylene ether glycol and at least one additional, but other glycol, selected from the group consisting of glycols and polyglycols.
- 10. Binder composition comprising a hydroxyl group containing polymer and an isocyanate crosslinking agent blocked with a blocking agent, characterized in that the blocking agent is an agent having a polymerizable double bond and an oxime-function.
- 11. Powder paint composition comprising a hydroxyl group containing polymer and an isocyanate crosslinking agent blocked with a blocking agent, characterized in that the blocking agent is an agent having a polymerizable double bond and an oxime-function.
- 12. An article coated with a composition according to any one of claims 1-4 or an article coated with a composition obtained according to any one of claims 6-8.

INTERNATIONAL SEARCH REPORT

Inten nal Application No
PCT/NL 93/00212

A. CLASSI	FICATION OF SUBJEC	T MATTER				
IPC 5	C08G18/12 C08G18/48	C08G18/28 C08G18/67	C08G18/62 C08G18/81	C08G18/66 C09D5/03		18/40 175/16
According to	o International Patent Cla	ssification (IPC) or to t	ooth national classificat	on and IPC		
B. FIELDS	SEARCHED					
Minimum d	ocumentation searched (o COSG	dassification system foll	owed by classification	rymbols)		
Documentat	ion searched other than n	ninimum documentation	to the extent that such	documents are included in	the fields se	arched
Electronic d	ata base consulted during	the international search	(name of data base ar	d, where practical, search t	erms used)	
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C. DOCUM	IENTS CONSIDERED T	O BE RELEVANT				
Category *	Citation of document, w	rith indication, where ag	propriate, of the releva	nt passages		Relevant to claim No.
A	FR,A,2 568 1986	884 (STEVENS	-GENIN) 14 F	ebruary		. 1
	see claims see page 1	1-3 , line 1 - li , line 16 - l	ne 4 ine 27			
A	EP,A,O 327 see claims	031 (ASAHI K 1,7,9	ASEI) 9 Augu 	st 1989		1
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Furt	her documents are listed i	n the continuation of bo	ox C.	Patent family member	s are listed in	n annex.
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Citation O' docume other r	n or other special reason (ent referring to an oral di neans	(as specified) sclosure, use, exhibition	or	cannot be considered to in document is combined wit ments, such combination in the art.	avolve an im th one or mo	ventive step when the are other such docu-
	int published prior to the nan the priority date clain		*&*	document member of the	same patent	family
Date of the	actual completion of the	international search		Date of mailing of the inte		rch report
4	February 1994	•		2 4. 62.	דנ	
Name and n		ice, P.B. 5818 Patentias	un 2	Authorized officer		
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Information on patent family members

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